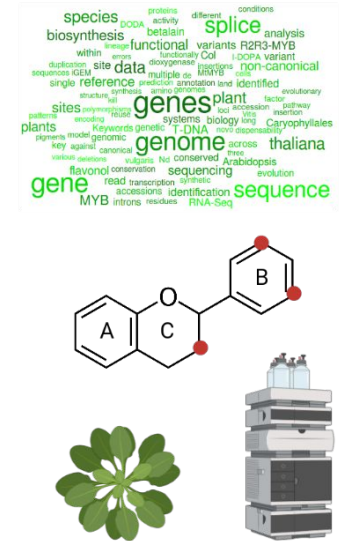
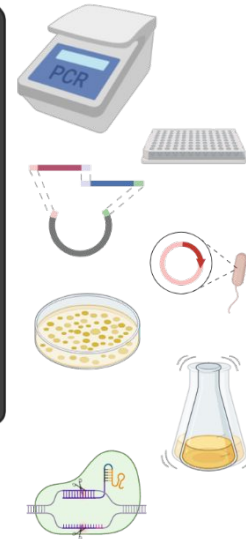
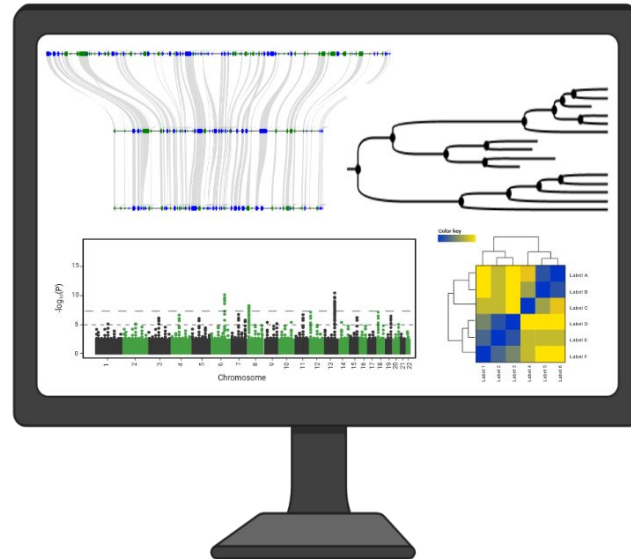
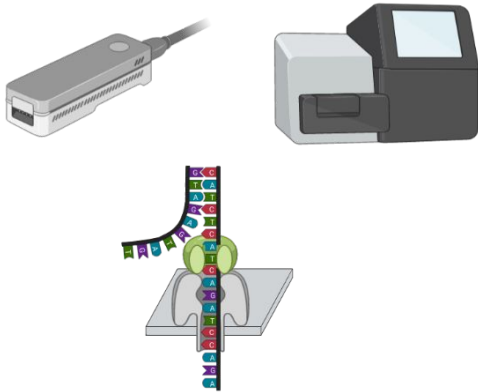




Technische
Universität
Braunschweig



Plant Biotechnology
and Bioinformatics

Python - File handling

Prof. Dr. Boas Pucker (Plant Biotechnology and Bioinformatics)

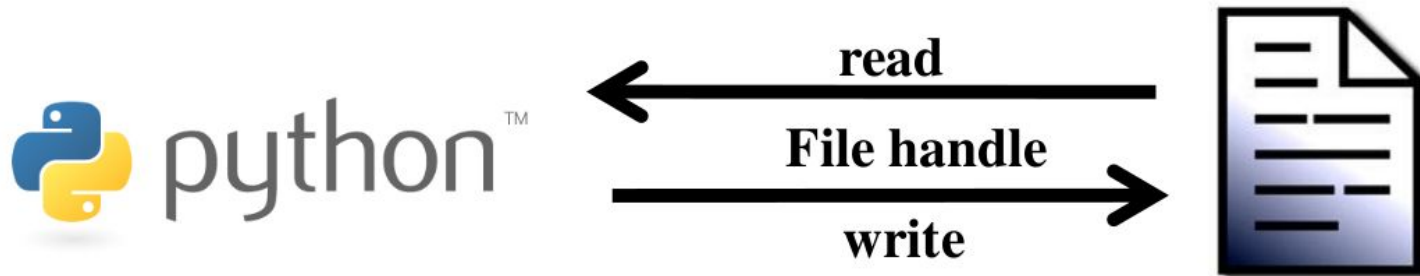
Availability of slides

- All materials are freely available (CC BY) - after the lectures:
 - StudIP: 'Python for Life Scientists'
 - GitHub: <https://github.com/bpucker/teaching>
- Questions: Feel free to ask at any time
- Feedback, comments, or questions: [b.pucker\[a\]tu-bs.de](mailto:b.pucker[a]tu-bs.de)

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Concept of file handling

- 'Connection' from Python to file
- Read = transfer of data from file
- Write = transfer of data into file



Reading a file (parsing)

```
f = open("test.txt", "r") #this opens a connection between Python and file
#f = connection from Python to file (file handle)
#'r' (reading) is default
lines = f.readlines() #reads content of file into a list
#each line becomes a list element of 'lines'
f.close() #close connection between Python and file
```

```
with open("test.txt", "r") as f: #opens file to read
    lines = f.readlines()
```

```
with open("test.txt", "r") as f: #opens file to read
    content = f.read() #reads entire file content into one string
```

Reading a file (big data)

- Advantage: only one line is read and processed at a time
- NGS data (e.g. FASTQ/SAM/BAM/VCF) are usually several GB in size
 - might exceed RAM
- Very long sequence (e.g. genome sequences) in FASTA
 - might be too large for available RAM

```
with open("text.txt", "r") as f: #"r" is default  
    #f = connection from Python to file (file handle)  
    line = f.readline() #reads next line  
    while line: #until end of file is reached  
        print(line)  
        line = f.readline() #reads next line
```

File analysis - example

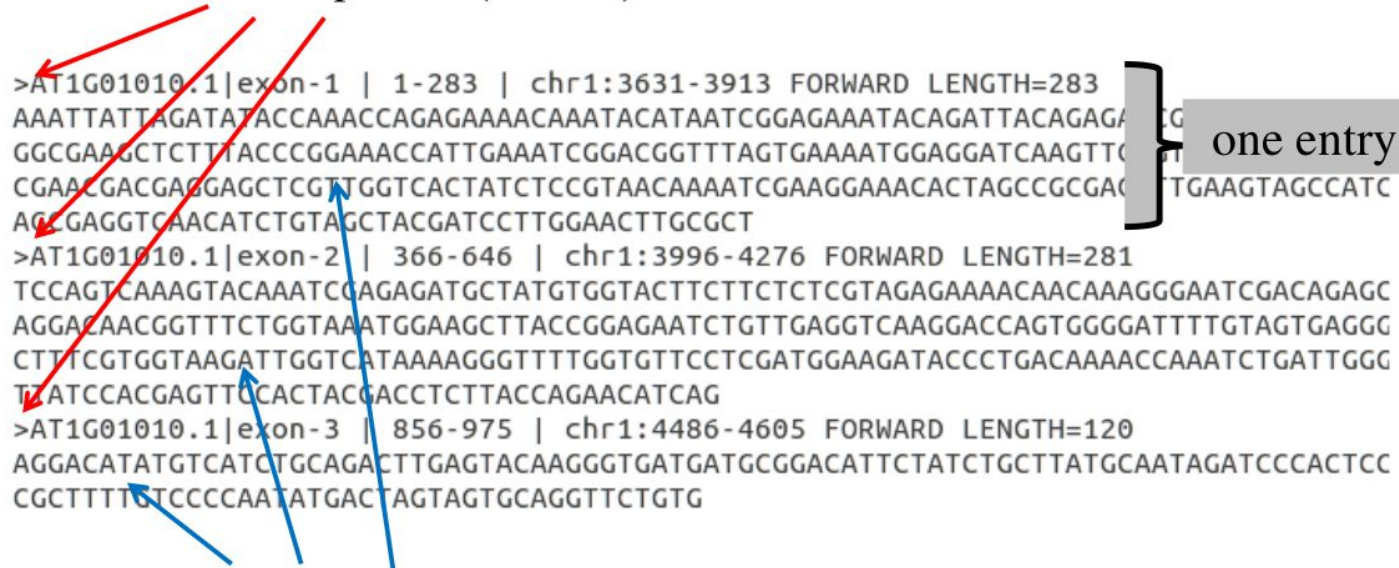
- How many lines are in AtCol0_Exons.fasta?
- Linux function: `$ head <FILENAME>`

```
>AT1G01010.1|exon-1 | 1-283 | chr1:3631-3913 FORWARD LENGTH=283
AAATTATTAGATATACCAAACCAGAGAAAACAAATACATAATCGGAGAAATACAGATTACAGAGAGCGAGAGAGATCGAC
GGCGAAGCTCTTTACCCGGAACCATTTGAAATCGGACGGTTTAGTGAAAATGGAGGATCAAGTTGGGTTTGGGTTCCGTC
CGAACGACGAGGAGCTCGTTGGTCACTATCTCCGTAACAAAATCGAAGGAAACACTAGCCGCGACGTTGAAGTAGCCATC
AGCGAGGTCAACATCTGTAGCTACGATCCTTGGAACCTGCGCT
>AT1G01010.1|exon-2 | 366-646 | chr1:3996-4276 FORWARD LENGTH=281
TCCAGTCAAAGTACAAATCGAGAGATGCTATGTGGTACTTCTTCTCTCGTAGAGAAAACAACAAAGGGAATCGACAGAGC
AGGACAACGGTTTCTGGTAAATGGAAGCTTACCGGAGAATCTGTTGAGGTCAAGGACCAGTGGGGATTTTGTAGTGAGGG
CTTTCGTGGTAAGATTGGTCATAAAAGGGTTTTGGTGTTTCCTCGATGGAAGATACCCTGACAAAACCAAATCTGATTGGG
TTATCCACGAGTTCCACTACGACCTCTTACCAGAACATCAG
>AT1G01010.1|exon-3 | 856-975 | chr1:4486-4605 FORWARD LENGTH=120
AGGACATATGTCATCTGCAGACTTGAGTACAAGGGTGATGATGCGGACATTCTATCTGCTTATGCAATAGATCCCACTCC
CGCTTTTGTCCCCAATATGACTAGTAGTGACAGGTTCTGTG
```


Multiple FASTA file

Name of sequence (header): line starts with '>'

```
>AT1G01010.1|exon-1 | 1-283 | chr1:3631-3913 FORWARD LENGTH=283  
AAATTATTAGATATACCAAACCAGAGAAAACAAATACATAATCGGAGAAATACAGATTACAGAGA  
GGCGAAGCTCTTTACCCGGAACCATTGAAATCGGACGGTTTAGTGAAAATGGAGGATCAAGTT  
CGAALGACGAGGAGCTCGTTGGTCACTATCTCCGTAACAAAATCGAAGGAAACACTAGCCGCGA  
AGCGAGGTCAACATCTGTAGCTACGATCCTTGGAACCTGCGCT  
>AT1G01010.1|exon-2 | 366-646 | chr1:3996-4276 FORWARD LENGTH=281  
TCCAGTCAAAGTACAAATCCAGAGATGCTATGTGGTACTTCTTCTCTCGTAGAGAAAACAACAAAGGGAATCGACAGAGC  
AGGACAACGGTTTCTGGTAAATGGAAGCTTACCGGAGAATCTGTTGAGGTCAAGGACCAGTGGGGATTTTGTAGTGAGGG  
CTTTCGTGGTAAGATTGGTCATAAAAGGGTTTTGGTGTTCTCGATGGAAGATACCCTGACAAAACCAAATCTGATTGGG  
TATCCACGAGTTCCACTACGACCTCTTACCAGAACATCAG  
>AT1G01010.1|exon-3 | 856-975 | chr1:4486-4605 FORWARD LENGTH=120  
AGGACATATGTCATCTGCAGACTTGAGTACAAGGTGATGATGCGGACATTCTATCTGCTTATGCAATAGATCCCACTCC  
CGCTTTTCTCCCAATATGACTAGTAGTGCAGGTTCTGTG
```



one entry

Sequence lines (no limit!)

Analyze FASTA file - example

```
1 with open( "/vol/apbiokurs/data/AtCol0_Exons.fasta", "r" ) as f:
2     line = f.readline() #reading first line
3     line_counter = 0 #counting lines
4     while line:
5         line_counter += 1 #counting lines
6         line = f.readline()
7     #number in line_counter needs to be converted to string:
8     print("File contains " + str( line_counter ) + " lines")
```


Exercises - Part4a

- 4.1) Count number of sequences (=number of headers) in example FASTA file!
- 4.2) Count number of sequence lines!
- 4.3) Count number of characters in document! (How many per line?)
- 4.4) How long are all contained sequences combined?
- 4.5) Calculate the average sequence length in this file!

And back again ... writing into file!

- If output file does not exist, it will be created
- File handle (f and out) can have any other name

```
1 Read:
2 with open( "test.txt", "r" ) as f:  #"r" (read) ist default
3     lines = f.readlines()
4
5
6
7
8 Write:
9 with open( "test2.txt", "w" ) as out:
10     out.write( "hello world!" )
```

difference: r = read; w = write

Writes a string into a file

Read & write

```
1 with open( "test.txt", "r" ) as f: #open file to read
2     with open( "test2.txt", "w" ) as out: #open file to write
3         line = f.readline() #read from first file
4         while line:
5             #this would be the place to apply filters
6             out.write( line )
7             line = f.readline()
```

Exercises - Part4b

- 4.6) Read the file AtCol0_Exons.fasta and write all headers (starting with '>') into a new file!
- 4.7) Read the file AtCol0_Exons.fasta and write the following into the output file:
 - The line if it is a header
 - The length of the line if it is a sequence line

White space characters

- New line (`'\n'`) and tab (`'\t'`) are special characters
 - Print `'/hello\tworld!\nhello\tworld!\n'`
- Python interprets these characters in print statements, but functions like `readline()` and `write()` do not!
 - New line (`'\n'`) needs to be added “manually” to each line

```
1 with open( "test_file.txt", "w" ) as out:  
2     out.write( "first test" )  
3     out.write( "second test" )  
4     out.write( "third test\n" )  
5     out.write( "fourth test\n" )  
6     out.write( "fifth test" )  
7     out.write( "sixth test" )
```

How many lines does
this file have?

strip()

- Removes white space characters from borders of a string
- often used to remove new lines ('\\n') at the line end

```
1 line = ">name_of_first_seq\\n"
2 print(line )
3 #>name_of_first_seq
4 # [empty line generated by \\n ]
5 line = line.strip()
6 print(line )
7 #>name_of_first_seq
```


split()

- Separates a string at each given occurrence of given substring
- Frequent separators: tab, comma, ...
- Generates list of strings

```
#tab-delimited file  
line = "column1\tcolumn2\tcolumn3\tcolumn4\tcolumn5\n"  
#line should be splitted at tabs  
columns = line.strip().split('\t')  
print(columns)  
#["column1", "column2", "column3", "column4", "column5"]
```

join()

- Combines strings of a list by putting a given substring between them
 - Examples: tab, space, comma, underscore
- Important: all elements of the list need to be string!

```
#tab-delimited file
line = "column1\tcolumn2\tcolumn3\tcolumn4\tcolumn5\n"
#line should be splitted at tabs
columns = line.strip().split('\t')
print(columns)
#[ "column1", "column2", "column3", "column4", "column5" ]


new_line = "_".join(columns)
print(new_line)
#column1_column2_column3_column4_column5
```

Exercises - Part4c

- 4.8) Calculate the number of sequences, the cumulative length, and the average length in the new file! Are they matching the values of the original file?
- 4.9) Write sequences into a new file if their length is a multiple of 10!
- 4.10) How many genes are located on Chr3?
- 4.11) What is the gene density of each chromosome (genes per Mbp)?
- 4.12) Write all sequences located on Chr2 between 10 and 15 Mbp.

Exercises - Part4d

- X4.1) Print the GC content of all genes with more than 10 exons.
- X4.2) Write the number of exons per gene into a new file.
- X4.3) Read the file AtCol0_Exons.fasta and write the following:
 - Only the Arabidopsis Gene Identifier (AGI, e.g. AT1G01010)
 - Gene identifier, exon name, and exon length (tab-delimited)



```
>AT1G01010.1|exon-1 | 1-283 | chr1:3631-3913 FORWARD LENGTH=283
AAATTATTAGATATACCAAACCAGAGAAAAACAAATACATAATCGGAGAAATACAGATTACAGAGAGCGAGAGAGATCGAC
GGCGAAGCTCTTTACCCGGAACCATTTGAAATCGGACGGTTTAGTGAAAATGGAGGATCAAGTTGGGTTTGGGTTCCGTC
CGAACGACGAGGAGCTCGTTGGTCACTATCTCCGTAACAAAATCGAAGGAAACACTAGCCGCGACGTTGAAGTAGCCATC
AGCGAGGTCAACATCTGTAGCTACGATCCTTGGAACCTTGCGCT
>AT1G01010.1|exon-2 | 366-646 | chr1:3996-4276 FORWARD LENGTH=281
TCCAGTCAAAGTACAAATCGAGAGATGCTATGTGGTACTTCTTCTCTCGTAGAGAAAACAACAAAGGGAATCGACAGAGC
AGGACAACGGTTTCTGGTAAATGGAAGCTTACCGGAGAATCTGTTGAGGTCAAGGACCAGTGGGGATTTTGTAGTGAGGC
CTTTCGTGGTAAGATTGGTCATAAAAGGGTTTTGGTGTTCCTCGATGGAAGATACCCTGACAAAACCAAATCTGATTGGC
TTATCCACGAGTTCCACTACGACCTCTTACCAGAACATCAG
>AT1G01010.1|exon-3 | 856-975 | chr1:4486-4605 FORWARD LENGTH=120
AGGACATATGTCATCTGCAGACTTGAGTACAAGGGTGATGATGCGGACATTCTATCTGCTTATGCAATAGATCCCACTCC
CGCTTTTGTCCCAATATGACTAGTAGTGCAGGTTCTGTG
```

Time for questions!